

Package ‘fanovaGraph’

October 7, 2020

Type Package

Title Building Kriging Models from FANOVA Graphs

Version 1.5

Date 2020-10-01

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Description Estimation and plotting of a function's FANOVA graph to identify the interaction structure and fitting, prediction and simulation of a Kriging model modified by the identified structure. The interactive function `plotManipulate()` can only be run on the 'RStudio IDE' with 'RStudio' package 'manipulate' loaded. 'RStudio' is freely available (<<https://rstudio.com/>>), and includes package 'manipulate'. The equivalent function `plotTk()` bases on CRAN Repository packages only. For further information on the method see Fruth, J., Roustant, O., Kuhnt, S. (2014) <doi:10.1016/j.jspi.2013.11.007>.

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LazyLoad yes

Depends sensitivity, igraph, DiceKriging (>= 1.4)

Suggests manipulate, testthat

NeedsCompilation no

Repository CRAN

Date/Publication 2020-10-07 12:10:02 UTC

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fanovaGraph-package *Building Kriging models with FANOVA graphs*

Description

Estimates and plots the FANOVA graph of a function to identify its interaction structure and fits a kriging model modified by the identified structure

Details

Important functions:

<code>estimateGraph</code>	Estimate indices for the graph, create graph structure
<code>threshold</code>	Set indices below a threshold to zero
<code>plot.graphlist</code>	Plot a given graph structure
<code>plotDeltaJumps</code>	Provide plots for the choice of the threshold
<code>kmAdditive</code>	Kriging model estimation with block-additive kernel
<code>predictAdditive</code>	Prediction function from Kriging model with block-additive kernel
<code>simAdditive</code>	Simulation from Kriging model with block-additive kernel

Author(s)

J. Fruth, T. Muehlenstaedt, O. Roustant, M. Jastrow

References

- Fruth, J.; Roustant, O.; Kuhnt, S. (2013+) Total interaction index: A variance-based sensitivity index for second-order interaction screening.
- Janon, A.; Klein, T.; Lagnoux-Renaudie, A.; Nodet, M.; Prieur, C. (2012+) Asymptotic normality and efficiency of two Sobol index estimators.
- Liu, R.; Owen, A.B. (2006) Estimating mean dimensionality of analysis of variance decompositions, *Journal of the American Statistical Association*, **101** 474, 712-721.
- Mara, T.A (2009) Extension of the RBD-FAST method to the computation of global sensitivity indices, *Reliability Engineering & System Safety*, **94** no. 8, 1274-1281.
- Muehlenstaedt, T.; Roustant, O.; Carraro, L.; Kuhnt, S. (2011) Data-driven Kriging models based on FANOVA-decomposition, *Statistics and Computing*.
- Sobol', I. M. (1993) Sensitivity estimates for nonlinear mathematical models, *Mathematical Modeling and Computational Experiment*, **1**, 407-414.

See Also

[DiceKriging](#), [sensitivity](#), [igraph](#)

Examples

```
#demo(ExampleIshigami)
#demo(Example6D)
#demo(Estimation)
#demo(Threshold)
```

estimateGraph	<i>FANOVA graph estimation.</i>
---------------	---------------------------------

Description

Estimates the structure of the FANOVA graph by estimating the total interaction indices for the graph edges (a particular case of superset importance introduced by Liu and Owen, 2006), the main effect indices for the graph vertices and the overall variance for normalizing the indices and finding the clique structure of the estimated graph.

Usage

```
estimateGraph(f.mat, d, q = NULL, q.arg = NULL, n.tot = NULL, method = "LiuOwen",
n.lo = NULL, n.mc = NULL, n.fast = 500, L = NULL, M = 6, n.pf = NULL, n.main = 1000,
confint = TRUE, print.loop.index = FALSE, ...)
```

Arguments

f.mat	vectorized function for which the FANOVA graph shall be estimated
d	integer, number of input factors (vertices)
q	a vector of character strings of quantile functions corresponding to the factors distributions, it can be a single character string meaning same distribution for all, if not specified "qunif" is taken
q.arg	a list of lists of quantile functions parameters of the distributions in q, it can be a single list meaning same parameters for all, if not specified the default values of the respective distributions are taken
n.tot	optional integer, total number of function evaluations, instead of n.tot method related parameters (n.lo, n.mc, L or n.sobol) can be provided
method	character string specifying the estimation method of the total interaction indices, to be chosen between "LiuOwen", "FixFast", "RBD" and "PickFreeze", defaults to "LiuOwen", see references for further details
n.lo	optional integer, only if method="LiuOwen", number of Monte Carlo simulations in method of Liu and Owen
n.mc	optional integer, only if method="FixFast", number of Monte Carlo simulations for the expectation in fixing method using FAST

n.fast	optional integer, only if method="FixFast", number of design points for FAST algorithm, defaults to 500
L	optional integer, only if method="RBD", parameter L in RBD-FAST method
M	optional integer, only if method="RBD", parameter M in RBD-FAST method
n.pf	optional integer, only if method="PickFreeze", number of Monte Carlo simulations in pick-and-freeze method
n.main	integer, number of Monte Carlo Simulations for computing main effect indices
confint	optional Boolean, if TRUE, standard error and 95% confidence intervals of the indices are computed additionally for method="LiuOwen", defaults to TRUE
print.loop.index	optional Boolean, if TRUE, loop indices are printed
...	additional arguments to be passed to the function f.mat

Value

an object of class `graphlist` containing the graph structure which includes

d	number of input factors
tii	matrix containing the unscaled total interaction indices and if <code>confint = TRUE</code> their standard error and lower and upper confidence limits
i1	matrix containing the unscaled main effect indices
V	overall variance
tii.scaled	matrix containing the scaled total interaction indices
cliques	list of cliques

Author(s)

J. Fruth, T. Muehlenstaedt

References

- Fruth, J.; Roustant, O.; Kuhnt, S. (2013+) Total interaction index: A variance-based sensitivity index for second-order interaction screening.
- Janon, A.; Klein, T.; Lagnoux, A.; Nodet, M.; Prieur, C. (2013) Asymptotic normality and efficiency of two Sobol index estimators.
- Liu, R.; Owen, A.B. (2006) Estimating mean dimensionality of analysis of variance decompositions, *Journal of the American Statistical Association*, **101** 474, 712-721.
- Mara, T.A (2009) Extension of the RBD-FAST method to the computation of global sensitivity indices, *Reliability Engineering & System Safety*, **94** no. 8, 1274-1281.
- Muehlenstaedt, T.; Roustant, O.; Carraro, L.; Kuhnt, S. (2011) Data-driven Kriging models based on FANOVA-decomposition, *Statistics and Computing*.
- Sobol', I. M. (1993) Sensitivity estimates for nonlinear mathematical models, *Mathematical Modeling and Computational Experiment*, **1**, 407-414.

Examples

```
# Ishigami function, true analytical values: D12 = D23 = 0, D13 =~ 3.374
q.arg = list(list(min=-pi, max=pi), list(min=-pi, max=pi), list(min=-pi, max=pi))
estimateGraph(f.mat=ishigami.fun, d=3, q.arg=q.arg, n.tot=10000, method="LiuOwen")
estimateGraph(f.mat=ishigami.fun, d=3, q.arg=q.arg, n.tot=10000, method="FixFast")
estimateGraph(f.mat=ishigami.fun, d=3, q.arg=q.arg, n.tot=10000, method="RBD")
estimateGraph(f.mat=ishigami.fun, d=3, q.arg=q.arg, n.tot=10000, method="PickFreeze")
```

i2Index

*Estimation of pure second order indices***Description**

Estimation of the unscaled pure second order Sobol indices.

Usage

```
i2Index(f.mat, d, q = NULL, q.arg = NULL, n.i2, ...)
```

Arguments

f.mat	vectorized function of which indices shall be estimated
d	integer, number of input factors (vertices)
q	a vector of character strings of quantile functions corresponding to the factors distributions, it can be a single character string meaning same distribution for all, if not specified "qunif" is taken
q.arg	a list of lists of quantile functions parameters of the distributions in q, it can be a single list meaning same parameters for all, if not specified the default values of the respective distributions are taken
n.i2	number of Monte Carlo evaluations
...	additional arguments to be passed to the function f.mat

Value

A vector containing the unscaled pure second order indices

Author(s)

J. Fruth

References

Sobol', I. M. (1993) Sensitivity estimates for nonlinear mathematical models, *Mathematical Modelling and Computational Experiment*, **1**, 407-414.

See Also

[estimateGraph totalIndex](#)

Examples

```
i2Index(f.mat=ishigami.fun, d=3, q.arg=list(min=-pi,max=pi), n.i2=10000)
```

 kmAdditive

Constrained MLE Optimization

Description

Constrained MLE optimization for kernels defined by cliques using [constrOptim](#)

Usage

```
kmAdditive(x, y, n.initial.tries = 50, limits = NULL, eps.R = 1e-08, cl,
covtype = "gauss", eps.Var = 1e-06, max.it = 1000, iso = FALSE)
```

Arguments

x	a design matrix of input variables, number of columns should be number of variables
y	a vector of output variables of the same length as the columns of x
n.initial.tries	number of random initial parameters for optimization, defaults to 50
limits	a list with items lower, upper containing boundaries for the covariance parameter vector theta, if NULL suitable bounds are computed from the range of x
eps.R	small positive number indicating the nugget effect added to the covariance matrix diagonalk, defaults to eps.R = 1e-08
cl	list of cliques, can be obtained by function threshold
covtype	an optional character string specifying the covariance structure to be used, to be chosen between "gauss", "matern5_2", "matern3_2", "exp" or "powexp" (see DiceKriging), defaults to "gauss"
eps.Var	small positive number providing the limits for the alpha parameters in order to guarantee strict inequalities ($0 + \text{eps.Var} \leq \alpha \leq 1 - \text{eps.Var}$), defaults to eps.Var = 1e-06
max.it	maximum number of iterations for optimization, defaults to max.it=1000
iso	boolean vector indicating for each clique if it is isotropic (TRUE) or anisotropic (FALSE), defaults to iso = FALSE (all cliques anisotropic)

Value

list of estimated parameter 'alpha' and 'theta' corresponding to the clique structure in 'cl'

Author(s)

T. Muehlenstadt, O. Roustant, J. Fruth

References

Muehlenstaedt, T.; Roustant, O.; Carraro, L.; Kuhnt, S. (2011) Data-driven Kriging models based on FANOVA-decomposition, *Statistics and Computing*.

See Also

[predictAdditive](#)

Examples

```
### example for ishigami function with cliques {1,3} and {2}
d <- 3
x <- matrix(runif(100*d, -pi, pi), nc=d)
y <- ishigami.fun(x)

cl <- list(c(2), c(1,3))

# constrained ML optimization with kernel defined by the cliques
parameter <- kmAdditive(x, y, cl = cl)

# prediction with the new model
xpred <- matrix(runif(500 * d, -pi, pi), ncol = d)
ypred <- predictAdditive(xpred, x, y, parameter, cl=cl)
yexact <- ishigami.fun(xpred)

# rmse
sqrt(mean((ypred[,1]- yexact)^2))

# scatterplot
par(mfrow=c(1,1))
plot(yexact, ypred[,1], asp = 1)
abline(0, 1)

### compare to one single clique {1,2,3}
cl <- list(c(1,2,3))

# constrained ML optimization with kernel defined by the cliques
parameter <- kmAdditive(x, y, cl = cl)

# prediction with the new model
ypred <- predictAdditive(xpred, x, y, parameter, cl=cl)

# rmse
sqrt(mean((ypred$mean- yexact)^2))

# scatterplot
par(mfrow=c(1,1))
plot(yexact, ypred$mean, asp = 1)
```

```
abline(0, 1)

### isotropic cliques

cl <- list(c(2),c(1,3))
parameter <- kmAdditive(x, y, cl = cl, iso=c(FALSE,TRUE))
ypred <- predictAdditive(xpred, x, y, parameter, cl=cl, iso=c(FALSE,TRUE))
sqrt(mean((ypred$mean- yexact)^2))

# the same since first clique has length 1
parameter <- kmAdditive(x, y, cl = cl, iso=c(TRUE,TRUE))
ypred <- predictAdditive(xpred, x, y, parameter, cl=cl, iso=c(TRUE,TRUE))
sqrt(mean((ypred$mean- yexact)^2))
```

kmPredictWrapper

Wrapper for the Kriging model prediction

Description

Wrapper for the Kriging model prediction function [predict.km](#) from package [DiceKriging](#) to simplify the use of Kriging prediction functions as arguments for functions like [estimateGraph](#) or [fast99](#).

Usage

```
kmPredictWrapper(newdata, km.object)
```

Arguments

newdata	a vector, matrix or data frame containing the points where to perform predictions
km.object	an object of class km

Value

kriging mean computed at newdata

Author(s)

J. Fruth, O. Roustant

See Also

[estimateGraph](#)

Examples

```

### graph estimation of a kriging prediction of the ishigami function
set.seed(1)
x <- matrix(runif(150,-pi,pi),100,3)
y <- ishigami.fun(x)
KM <- km(~1, design = data.frame(x), response = y)

g <- estimateGraph(f.mat = kmPredictWrapper, d = 3, n.tot = 10000, q.arg =
  list(min = -pi, max = pi), method = "LiuOwen", km.object = KM)
print(g$tii)

```

L

*LHS Dataset***Description**

6-dimensional Latin Hypercube Sampling Dataset

Usage

```
data(L)
```

Format

The format is: num [1:100, 1:6] -0.7105 -0.7739 -0.5017 0.6158 0.0245 ...

Examples

```
data(L)
## str(L) ; pairs(L) ...
```

plot.graphlist

*Plot Graph via Package [igraph](#)***Description**

Plot FANOVA graphs using functions from package [igraph](#).

Usage

```

## S3 method for class 'graphlist'
plot(x, names = NULL, i2 = NULL, layout = NULL, plot.i1=TRUE, max.thickness=15,
circle.diameter=40, ...)

```

Arguments

x	an object of class <code>graphlist</code> as obtained from estimateGraph
names	optional character string, names of vertices, defaults to 1:d
i2	optional vector of second order interaction indices (thickness of inner edges)
plot.i1	optional boolean, if TRUE main effects are drawn in the graph by vertices thicknesses, should be set to FALSE when only total interaction indices are of interest
layout	optional layout for the graph as in igraph , default is <code>layout.fruchterman.reingold</code>
max.thickness	optional value for the maximal line thickness, defaults to 20
circle.diameter	optional value for the circle diameter, defaults to 40
...	additional arguments, passed to plot

Author(s)

J. Fruth, O. Roustant, S. Hess, S. Neumaerker

References

Muehlenstaedt, T.; Roustant, O.; Carraro, L.; Kuhnt, S. (2011) Data-driven Kriging models based on FANOVA-decomposition, *Statistics and Computing*.

Csardi, G.; Nepusz, T. (2006) The igraph software package for complex network research, *Inter-Journal Complex Systems*, **Complex Systems**, 1695.

See Also

[plotGraphChange](#)

Examples

```
op <- par(no.readonly=TRUE)
g1 <- estimateGraph(f.mat=ishigami.fun, d=3, q.arg=list(min=-pi,max=pi), n.tot=10000)
plot(g1)
plot(g1, names=c("A","B","C"))
plot(g1, names=c("A","B","C"), plot.i1 = FALSE)

# include pure second order indices
g2 <- estimateGraph(f.mat=function(x) x[,1]*x[,2]*x[,3]+x[,2]*x[,3], d=3,
  q.arg=list(min=-1,max=1), n.tot=10000)

plot(g2)
plot(g2, i2 = c(0.001, 0.001, 0.05))

# equal layouts and different edge thicknesses and circle diameters
g3 <- estimateGraph(f.mat=function(x) x[,1]*x[,2]*x[,3]*x[,4]*x[,5], d=5,
  q.arg=list(min=-1,max=1), n.tot=10000)

g4 <- estimateGraph(f.mat=function(x) x[,1]*x[,2]*x[,3]+x[,4]*x[,5], d=5,
  q.arg=list(min=-1,max=1), n.tot=10000)
```

```
graphClassIgraph <- graph.full(n = 5, directed = FALSE)
layout <- layout.circle(graphClassIgraph)

plot(g3, max.thickness= 10, circle.diameter= 30, layout=layout)
plot(g4, max.thickness= 30, circle.diameter= 50, layout=layout)
```

plotDeltaJumps	<i>Delta Jump Plot</i>
----------------	------------------------

Description

Threshold discision plot. plotDeltaJumps plots the threshold steps (the values of delta at which the graph changes) equidistant against the number of cliques and the values of delta on the real axis. The indices are assumed to be scaled for the threshold cuts.

Usage

```
plotDeltaJumps(graphlist, interval = c(0, 1), mean.clique.size = FALSE)
```

Arguments

graphlist	an object of class graphlist as obtained from estimateGraph
interval	an optional vector of size 2, range for the values of delta to be shown in the plot, defaults to c(0,1)
mean.clique.size	logical, if TRUE (default) an additional line is drawn representing the mean of the number of vertices in the cliques

Details

The plots shall give help in the choice for the treshold. In the first plot a small number of cliques might be preferable in order to have less parameters to estimate. If several values result in the same number of cliques the ones with higher mean clique size are possibly preferable.

In the second plot a high jump indicates a point of big distance between two successive edge indices and thus a clear change in the graph structure. The intervals with notable jumps are highlighted in green, the higher the jump the darker the colour. Those highlighted intervals together with a small number of cliques are probably good choices for the threshold.

Use [plotGraphChange](#) to visualize the graph structure for possible threshold values.

Author(s)

J. Fruth, O. Roustant

See Also

[thresholdIdentification](#), [plotGraphChange](#)

Examples

```
tii <- matrix(c(0.0018, 0.0265, 0.0017, 0.0277, 0.0018, 0.001, 0.028, 0.0013,
  0.0212, 0.002, 0.0372, 0.0024, 0.0022, 0.0157, 0.003))
g <- list(d = 6,
  tii = tii,
  i1 = matrix(c(0.0901, 0.1288, 0.0683, 0.0979, 0.0882, 0.1572)),
  V = 0.8,
  tii.scaled = tii/0.8,
  cliques = list(1:6))

### Delta Jump Plot (jump between 0.0038 and 0.0196)
plotDeltaJumps(g)
```

plotGraphChange, plotTk, plotManipulate
Plot Graph as It Changes with Delta

Description

Graphs are plotted depending on a change on delta, the threshold for edges to appear in the graph, to enable a visual decision for delta by graph behavior.

Usage

```
plotGraphChange(graphlist, fix.layout = TRUE, delta.layout = 0.01)
plotTk(graphlist, delta.layout=0.01)
plotManipulate(graphlist, delta.layout=0.01)
```

Arguments

graphlist	an object of class graphlist as obtained from estimateGraph
fix.layout	logical, if TRUE (default) the position of the vertices is fixed for all plots such that the positions are optimal for delta = delta.layout
delta.layout	optional value between 0 and 1, see fix.layout, defaults to 0.01

Note

plotGraphChange shows the changing of the graph step by step by changing plots as in demo, plotTk is an interactive version using [tcltk](#), plotManipulate is an interactive version using manipulate

Author(s)

J. Fruth, O. Roustant

See Also

[plotDeltaJumps](#), [plot.graphlist](#)

Examples

```
# see demo(Threshold)
```

predictAdditive	<i>Prediction Function with Modified Kernel</i>
-----------------	---

Description

Standard kriging prediction function for the modified correlation functions.

Usage

```
predictAdditive(newdata, x, y, parameter, covtype = "gauss", eps.R = 1e-08,
  cl, iso = FALSE, se.compute=FALSE)
```

Arguments

newdata	matrix containing the points where to perform predictions
x	matrix of input data
y	vector of output data
parameter	(by kmAdditive estimated) kriging parameters, list of size of 'cl' containing for each clique a list of parameters alpha (single value) and theta (numeric vector of values)
covtype	an optional character string specifying the covariance structure to be used, to be chosen between "gauss", "matern5_2", "matern3_2", "exp" or "powexp" (see DiceKriging), defaults to "gauss"
eps.R	small positive number indicating the nugget effect added to the covariance matrix diagonalk, defaults to eps.R = 1e-08
cl	list of cliques
iso	boolean vector indicating for each clique if it is isotropic (TRUE) or anisotropic (FALSE), defaults to iso = FALSE (all cliques anisotropic)
se.compute	optional boolean. If FALSE, only the kriging mean is computed. If TRUE, the kriging variance (actually, the corresponding standard deviation) is computed, too

Value

mean	kriging mean computed at newdata.
sd	kriging standard deviation computed at newdata. Only computed if se.compute=TRUE.

Author(s)

T. Muehlenstaedt, O. Roustant, J. Fruth

References

Muehlenstaedt, T.; Roustant, O.; Carraro, L.; Kuhnt, S. (2011) Data-driven Kriging models based on FANOVA-decomposition, *Statistics and Computing*.

See Also

[kmAdditive](#)

Examples

```
### example for ishigami function with cliques {1,3} and {2}
d <- 3
x <- matrix(runif(100*d, -pi, pi), nc=d)
y <- ishigami.fun(x)

cl <- list(c(2), c(1,3))

# constrained ML optimization with kernel defined by the cliques
parameter <- kmAdditive(x, y, cl = cl)

# prediction with the new model
xpred <- matrix(runif(500 * d, -pi, pi), ncol = d)
ypred <- predictAdditive(xpred, x, y, parameter, cl=cl)
yexact <- ishigami.fun(xpred)

# rmse
sqrt(mean((ypred[,1]- yexact)^2))

# scatterplot
par(mfrow=c(1,1))
plot(yexact, ypred[,1], asp = 1)
abline(0, 1)

### compare to one single clique {1,2,3}
cl <- list(c(1,2,3))

# constrained ML optimization with kernel defined by the cliques
parameter <- kmAdditive(x, y, cl = cl)

# prediction with the new model
ypred <- predictAdditive(xpred, x, y, parameter, cl=cl)

# rmse
sqrt(mean((ypred$mean- yexact)^2))

# scatterplot
par(mfrow=c(1,1))
plot(yexact, ypred$mean, asp = 1)
```

```

abline(0, 1)

### isotropic cliques

cl <- list(c(2),c(1,3))
parameter <- kmAdditive(x, y, cl = cl, iso=c(FALSE,TRUE))
ypred <- predictAdditive(xpred, x, y, parameter, cl=cl, iso=c(FALSE,TRUE))
sqrt(mean((ypred$mean- yexact)^2))

# the same since first clique has length 1
parameter <- kmAdditive(x, y, cl = cl, iso=c(TRUE,TRUE))
ypred <- predictAdditive(xpred, x, y, parameter, cl=cl, iso=c(TRUE,TRUE))
sqrt(mean((ypred$mean- yexact)^2))

```

simAdditive

Simulate GP values from block-additive kernel

Description

Simulate Gaussian process values from a given block-additive kernel

Usage

```
simAdditive(newdata, mu, parameter, covtype, cl, iso = FALSE, eps.R = 1e-08)
```

Arguments

newdata	matrix containing the points where to perform simulations
mu	trend parameter
parameter	list of size of 'cl' containing for each clique a list of parameters alpha (single value) and theta (numeric vector of values)
covtype	character string specifying the covariance structure to be used, to be chosen between "gauss", "matern5_2", "matern3_2", "exp" or "powexp" (see DiceKriging)
cl	list of cliques
iso	boolean vector indicating for each clique if it is isotropic (TRUE) or anisotropic (FALSE), defaults to iso = FALSE (all cliques anisotropic)
eps.R	small positive number indicating the nugget effect added to the covariance matrix diagonalk, defaults to eps.R = 1e-08

Value

a vector containing the simulated values

Author(s)

J. Fruth

References

Muehlenstaedt, T.; Roustant, O.; Carraro, L.; Kuhnt, S. (2011) Data-driven Kriging models based on FANOVA-decomposition, *Statistics and Computing*.

Rasmussen, C. E.; Williams, C. K. I. (2006), *Gaussian processes for machine learning*, MIT Press.

See Also

[kmAdditive](#), [simulate](#)

Examples

```
### 2 dimensional simulation
x1 <- x2 <- seq(-1,1,,20)
x <- expand.grid(x1,x2)
covtype <- "matern3_2"
mu <- 0

op <- par(no.readonly=TRUE); par(mfrow=c(1,2), mar=c(1,1,1,1))
# non-additive simulation
parameter <- list(list(alpha=1, theta=c(0.8,0.8)))
c1 <- list(1:2)
set.seed(1)
y <- simAdditive(x, mu, parameter, covtype, c1)
persp(x1,x2, matrix(y,20), theta=-40, col="lightblue", zlab="y")

# additive simulation
parameter <- list(list(alpha=0.5, theta=0.8),
                  list(alpha=0.5, theta=0.8))
c1 <- list(1,2)
set.seed(1)
y <- simAdditive(x, mu, parameter, covtype, c1)
persp(x1,x2, matrix(y,20), theta=-40, col="lightblue", zlab="y")

par(op)
```

threshold

Threshold indices

Description

All indices below a treshold are set to be zero.

Usage

```
threshold(graphlist, delta, scaled = TRUE, robust = FALSE)
```


Arguments

graphlist	an object of class <code>graphlist</code> as obtained from <code>estimateGraph</code>
delta	numeric threshold, between 0 and 1 if <code>scaled = TRUE</code>
scaled	optional boolean, if <code>TRUE</code> , indices are normalized by the overall variance before for threshold cut, defaults to <code>TRUE</code>
robust	optional boolean, if <code>TRUE</code> , upper confidence intervals limits are used for the threshold cut instead of indices themselves, confidence intervals must be provided in <code>graphlist</code> , defaults to <code>FALSE</code>

Value

an object of class `graphlist` where the indices are thresholded the clique structure is updated respectively, see `estimateGraph` for a detailed description

Warning

The threshold cut is by default performed on scaled indices. For a cut on the original unscaled indices set `scaled = FALSE`.

Author(s)

J. Fruth, T. Muehlenstaedt, O. Roustant

References

Muehlenstaedt, T.; Roustant, O.; Carraro, L.; Kuhnt, S. (2011) Data-driven Kriging models based on FANOVA-decomposition, *Statistics and Computing*.

Examples

```
# Kriging model prediction
x <- matrix(runif(100*3,-pi,pi),100,3)
KM <- km(~1, design = data.frame(x), response = ishigami.fun(x))
krigingMean <- function(Xnew) predict(object = KM, newdata = Xnew,
  type = "UK", se.compute = FALSE, checkNames = FALSE)$mean

# full graph estimation
g <- estimateGraph(krigingMean, d=3, n.tot=10000, q.arg=list(min=-pi, max=pi))
print(g[c(2,6)])
# threshold graph
g.cut <- threshold(g, delta = 0.1)
print(g.cut[c(2,6)])
```

 thresholdIdentification

Function to identify a suitable threshold for an estimateGraph object

Description

From an estimateGraph object and a corresponding data set, candidate threshold values are compared on the prediction performance of the corresponding additive Kriging model. The candidate thresholds are chosen by the biggest jumps in plotDeltaJumps together with 0 (the full model) and 1 (the complete additive model). For each of them the Kriging model with corresponding kernel is estimated and the leave-one-out crossvalidations on the original data sets are compared on scatterplots and RMSE-values.

Usage

```
thresholdIdentification(g, x, y, n.cand = 3, covtype = "matern5_2", KM = NULL)
```

Arguments

g	object of class graphlist as obtained from estimateGraph
x	design matrix of input variables corresponding to g, number of columns should be number of variables
y	vector of output variables of the same length as the columns of x
n.cand	integer, the n.cand biggest jumps are chosen as candidate threshold values. The default value is 3
covtype	optional character string specifying the covariance structure to be used. The default is "matern5_2"
KM	optional object of class km, the full kriging model corresponding to g. With default value NULL, this kriging model is computed by the function itself

Value

a list including

delta	vector of threshold candidates
models	list of full model and models with applied thresholds
y.cv	list of vectors containing crossvalidation predictions for each model
RMSE	vector of residual mean squared errors for each model

Author(s)

J. Fruth, M. Jastrow

See Also

[plotDeltaJumps](#), [plotGraphChange](#)

Examples

```
##### simple 3-dimensional example with one interaction

### data (usually existing)
x <- matrix(seq(0,1,,20), 20, 3)
x <- apply(x,2,sample)
y <- 2*(x[,1]-0.5) * (x[,2]-0.5) + 0.1*sin(10*x[,3])

### FANVOA graph (usually estimated from a meta model over the data)
g <- list(d=3,
  tii = matrix(c(0.0140, 0.0008, 0.0002)),
  V = 0.0222,
  tii.scaled = matrix(c(0.6976, 0.0432, 0.0113))
)
class(g) <- "graphlist"

### plot complete graph
plot(g, plot.i1=FALSE)

### Compare candidate thresholds on prediction performance
set.seed(1)
comparison <- thresholdIdentification(g, x, y, n.cand = 1)
```

totalIndex

*Estimation of main index indices***Description**

Estimation of the unscaled total Sobol index of all main indices by method Liu & Owen (superset importance of main indices).

Usage

```
totalIndex(f.mat, d, q = NULL, q.arg = NULL, n.mc, ...)
```

Arguments

f.mat	vectorized function of which indices shall be estimated
d	integer, number of input factors (vertices)
q	a vector of character strings of quantile functions corresponding to the factors distributions, it can be a single character string meaning same distribution for all, if not specified "qunif" is taken
q.arg	a list of lists of quantile functions parameters of the distributions in q, it can be a single list meaning same parameters for all, if not specified the default values of the respective distributions are taken
n.mc	number of Monte Carlo evaluations
...	additional arguments to be passed to the function f.mat

Value

A vector containing the unscaled total Sobol indices

Author(s)

J. Fruth

References

Liu, R.; Owen, A.B. (2006) Estimating mean dimensionality of analysis of variance decompositions, *Journal of the American Statistical Association*, **101** 474, 712-721.

See Also

[estimateGraph](#)

Examples

```
totalIndex(f.mat=ishigami.fun, d=3, q.arg=list(min=-pi,max=pi), n.mc=10000)
totalIndex(f.mat=sobol.fun, d=8, q.arg=list(min=0,max=1), n.mc=10000)
```

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